

Michael R. Pence Governor

William C. VanNess II, MD State Health Commissioner

DATE:

September 3, 2014

TO:

All Local Health Departments

Attn: Chief Food Inspection Officer

FROM:

Laurie Kidwell, RRT Supervisor

Food Protection Program

SUBJECT:

Regeneca Worldwide a division of VivaCeuticals, Inc. – RECALL [Drug]

EXPANDED RECALL

AFFECTED

PRODUCT:

RegeneSlim appetite control dietary supplement

SUMMARY:

Unclassified Recall; The recall is due to FDA analysis that has confirmed the presence of DMAA. DMAA is also known as 1,3-dimethylamylamine, methylhexanamine, or geranium extract. The Food and Drug Administration (FDA) has warned that DMAA is potentially dangerous to health as it can narrow blood vessels and arteries, which can cause a rise in blood pressure or other cardiovascular problems such as shortness of

breath, arrhythmias, tightening in the chest, and heart attack.

RegeneSlim is packaged in approximately 3 ½" by 3" green and white sachets that contain 2 capsules, with the name RegeneSlim displayed prominently on the front of the sachet to include lot #823230415, lot #EX0616r 15813, Lot # EX0616R15814 and Lot

#11414re5516.

RegeneSlim is purchased by and distributed through a direct sales force within the <u>United States</u> and Puerto Rico, and through online sales, for both personal consumption

and retail sales.

SUGGESTED

ACTION:

For consumer inquiry only. Consumers with questions may contact the company at 1-949-281-2600 between the hours of 9 a.m. and 6 p.m. PDT. Consumers should contact their physician or healthcare provider if they experience any problems that may be related to taking or using RegeneSlim.

Recall -- Firm Press Release



FDA posts press releases and other notices of recalls and market withdrawals from the firms involved as a service to consumers, the media, and other interested parties. FDA does not endorse either the product or the company.

Regeneca Worldwide, A Division Of Vivaceuticals, Inc Voluntarily Recalls RegeneSlim Appetite Control Capsules Due To The Presence Of DMAA That May Pose Possible Health Risk

Contact

Consumer/Media: 949-281-2600

FOR IMMEDIATE RELEASE – August 6, 2014 – Regeneca Worldwide a division of VivaCeuticals, Inc. Las Vegas, NV is conducting a voluntary nationwide recall of its RegeneSlim appetite control dietary supplement from lot # EX0616R15814 and lot #11414RE5516 because FDA analysis confirmed the presence of DMAA. DMAA is also known as 1,3-dimethylamylamine, methylhexanamine, or geranium extract. DMAA is commonly used as a stimulant, pre-workout, and weight loss ingredient in dietary supplement products. The Food and Drug Administration (FDA) has warned that DMAA is potentially dangerous to health as it can narrow blood vessels and arteries, which can cause a rise in blood pressure or other cardiovascular problems such as shortness of breath, arrhythmias, tightening in the chest, and heart attack.

RegeneSlim is purchased by and distributed through a direct sales force within the United States and Puerto Rico, and through online sales, for both personal consumption and retail sales.

RegeneSlim is packaged in approximately 3 ½" by 3" green and white sachets that contain 2 capsules, with the name RegeneSlim displayed prominently on the front of the sachet.

There have been no illnesses reported to date.

This voluntary recall was the result of FDA analysis confirming the presence of DMAA in RegeneSlim and our company's sampling. The company continues their investigation as to what caused the problem.

Consumers who have purchased RegeneSlim with the above-mentioned lot numbers are advised to immediately stop using the product and are urged to return it to the place of purchase for a full exchange. Consumers with questions may contact the company at 1-949-281-2600 between the hours of 9 a.m. and 6 p.m. PDT. Consumers should contact their physician or healthcare provider if they experience any problems that may be related to taking or using RegeneSlim.

Adverse reactions or quality problems experienced with the use of this product may be reported to the FDA's MedWatch Adverse Event Reporting program either online, by regular mail or by fax.

Complete and submit the report Online: www.fda.gov/medwatch/report.htm

Regular Mail or Fax: Download form www.fda.gov/MedWatch/getforms.htm or call 1-800-332-1088 to request a reporting form, then complete and return to the address on the pre-addressed form, or submit by fax to 1-800-FDA-0178.

This recall is being conducted with the knowledge of the U.S. Food and Drug Administration.

Regeneca Worldwide, a Division of Vivaceuticals, Inc Expands the Voluntarily Recall of Regeneslim Appetite Control Capsules Due to the Presence of DMAA that May Pose Possible Health Risk Photos



RegeneSlim™ Appetite Control

Amount Per 2 Servings		% Dally Value*		
Chromium (as 0-polynicotinate) Garcinia Cambogia (fruit) Extract† (-) Hydroxycitric Acid 60%	375 mg	83,5%		
Calcium (as Hydroxycitrate) Potassium (as Hydroxycitrate)		7.5% 1.5%		

176 mg **Proprietary Blend** Kola Nut Extract 10%, Green Coffee Bean 50% Chlorogenic Acid, R-beta-Methylphenylethylamine HCL, HCL, Synephrine HCL 98%, Guggulsterones Natural 10%, Tyramine HCL 98%, Hamala Extract 40%, Coleus 10%.

- "Pero nt Dally Values are based on a 2,000 calorie diet.
- "Dail Value not established.
- † Super CitriMax® brand (-)hydroxycitric acid (U.S. Patents 6,160,172, 6,395,296, 6,875,891, 7,335,651 and worldwide patents pending). OTHER INGREDIENTS: Gelatin Capsule, Caffeine



CHROMEMATE®

+ Super CitriMax* and ChromeMate* are trademarks of InterHealth N.I.

DIRECTIONS FOR USE: Take 1 capsulg twice daily, preferably 1 to 2 hours before or between meals.

READ ENTIRE LABEL BEFORE TAKING.

Store at room temperature,

Do not use if seal is broken or missing.

KEEP OUT OF REACH OF CHILDREN.

WARNING: Not for use by individuals under 18. Do not use if pregnant or nursing. May contain pollen, which can cause reactions in some people. If rash or other symptoms appear, discontinue use and consult your healthcare professional. Do not use if you are at risk or are being treated for high blood pressure, kidney, thyrold, heart or psychiatric disease, anxiety, depression, seizure disorder or stroke. Individuals who consume caffeire with this product may experience adverse effects. Seek professional assistance in case of accidental overdose or if you experience repid heartbest, dizzness, severe headache or shortness of breath. Consult with your healthcare professional prior to use if you have a medical condition or are taking medication, if sleeplessness occurs discontinue affarmoon dose.

*These statements have not been evaluated by the Food and Drug Admira-This product is not intended to diagnose, treat, cure or prevent disease.



Olstributed By:

Regeneca Worldwide^{nt} 16 Technology Dr., Suite 124, Irvine, CA 92618 (949) 281-2600 | www.RegeneStim.com

Lot#: